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No. 718982

STANDARD PATENT

I, Vivienne Joyce Thom, Commissioner of Patents, grant a Standard Patent with the following particulars:

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- (54) Title: DNA SEQUENCE OF A GENE OF HYDROXY-PHENYL PYRUVATE DIOXYGENASE AND PRODUCTION OF PLANTS CONTAINING A GENE OF HYDROXY-PHENYL PYRUVATE DIOXYGENASE AND WHICH ARE TOLERANT TO CERTAIN HERBICIDES
- (54) Titre: SEQUENCE ADN D'UN GENE DE L'HYDROXY-PHENYL PYRUVATE DIOXYGENASE ET OBTENTION DE PLANTES CONTENANT UN GENE DE L'HYDROXY-PHENYL PYRUVATE DIOXYGENASE, TOLERANTES A CERTAINS HERBICIDES

(57) Abstract

DNA sequence of a gene of hydroxy-phenyl pyruvate dioxygenase and production of plants containing a gene of hydroxy-phenyl pyruvate dioxygenase and which are resistant to herbicides. DNA sequence of a gene of hydroxy-phenyl pyruvate dioxygenase; isolation from a bacteria or a plant; utilization for obtaining plants tolerant to herbicides.

(57) Abrégé

Séquence ADN d'un gène de l'hydroxy-phényl pyruvate dioxygénase et obtention de plantes contenant un gène de l'hydroxy-phényl pyruvate dioxygénase, résistantes aux herbicides. Séquence ADN d'un gène de l'hydroxy-phényl pyruvate dioxygénase; isolement à partir d'une bactérie ou d'une plante; utilisation pour l'obtention de plantes tolérantes aux herbicides.

WO 96/38567 PCT/FR96/00831

DNA sequence of a hydroxyphenylpyruvate dioxygenase gene and obtainment of plants comprising a hydroxyphenylpyruvate dioxygenase gene, which are tolerant to certain herbicides

The present invention relates to a hydroxyphenylpyruvate dioxygenase (HPPD) gene, a chimeric gene comprising this gene as coding sequence and its use to obtain plants resistant to certain herbicides.

Certain herbicides have been disclosed, such as the isoxazoles described especially in the French Patent Applications 95 06800 and 95 13570 and especially isoxaflutole, a selective maize herbicide, diketonitriles such as those described in European

Applications 0 496 630, 0 496 631, in particular 2-cyano-3-cyclopropyl-1-(2-SO₂CH₃-4-CF₃-phenyl)propane-1,3-dione and 2-cyano-3-cyclopropyl-1-(2-SO₂CH₃-4-2,3Cl₂phenyl)propane-1,3-dione, triketones described in European Applications 0 625 505 and 0 625 508, in particular sulcotrione. However, a tolerance gene to such herbicides has not been described.

Hydroxyphenylpyruvate dioxygenase is an enzyme which catalyses the conversion reaction of parahydroxyphenylpyruvate into homogentisate.

In addition, the amino-acid sequence of hydroxyphenylpyruvate dioxygenase from Pseudomonas sp. P.J. 874 has been described, without there being a

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description of its role in the tolerance of the plants to herbicides (Rüetschi et al.: Eur. J. Biochem. 205, 459-466, 1992). This document does not give a description of the gene coding for this protein.

of a gene of this type and that such a sequence could, once incorporated into plant cells, produce an over-expression or an activation of HPPD in the plants giving to the latter a worthwhile tolerance to certain novel herbicides, such as those of the isoxazoles family or that of the triketones.

The present invention accordingly provides an isolated sequence of a gene expressing a hydroxyphenylpyruvate dioxygenase (HPPD), characterized in that it is from Pseudomonas fluorescens.

The invention also provides an isolated sequence expressing a HPPD, characterized in that it is of plant origin, such as especially of monocotyledonous or dicotyledonous plants, especially of Arabidopsis or of Umbelliferae, such as, for 20 example, the carrot (Daucus carotta). It can be native or wild or possibly mutated while at the same time fundamentally retaining a property of herbicidal tolerance against HPPD inhibitors, such as herbicides of the isoxazoles family or that of the triketones. The sequences shown in SEQ ID NO:2 and SEQ 25 ID NO:3 are preferred.



The invention likewise relates to a process of isolating the above gene, characterized in that:

- as primers, some oligonucleotides from the amino-acid sequence of an HPPD are chosen,
- 5 starting from these primers, amplification fragments are synthesized by PCR
 - the gene is isolated by creation and screening of a genomic bank and
 - the gene is cloned.
- Preferably, primers from the HPPD sequence of a bacterium of the genus Pseudomonas are used.

 Particularly preferably, they are from Pseudomonas fluorescens.

The invention also relates to the use of a

15 gene coding for HPPD in a process for the
transformation of plants, as a marker gene or as a
coding sequence allowing tolerance to certain
herbicides to be conferred on the plant. It can
likewise be used, in association with other marker

20 genes and/or coding sequence, for an agronomic
property.

The coding gene can be of any origin, native or wild or possibly mutated, while at the same time fundamentally retaining a property of herbicidal

25 tolerance against inhibitors of HPPD, such as herbicides of the isoxazoles family or that of the triketones. As coding sequence, especially that according to the invention such as described above, can

be used.

The transformation of plant cells can be achieved by any appropriate known means. A series of methods consists in bombarding cells or protoplasts with particles to which are coupled the DNA sequences.

Another series of methods consists in using, as means of transfer into the plant, a chimeric gene inserted into a Ti plasmid of Agrobacterium tumefaciens or Ri plasmid of Agrobacterium rhizogenes.

An object of the present invention is also a chimeric 10 gene for the genetic transformation of plants comprising, in the transcription direction, at least one promoter regulation sequence from a gene expressed naturally in plants, a heterologous coding sequence (for example of bacterial, such as which expresses of plant origin) 15 Pseudomonas OT SD. one hydroxyphenylpyruvate dioxygenase and at least polyadenylation sequence.

The promoter regulation sequence used can be any promoter sequence of a gene which is naturally 20 expressed in plants, in particular a promoter of bacterial, viral or plant origin, such as, for example, that of a gene of the small subunit of ribulose biscarboxylase (RuBisCO) or that of a gene of α-tubulin (European Application EP No. 0 652 286), or 25 alternatively of a plant virus gene such as, for example, that of cauliflower mosaic virus (CAMV 195 or



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35S), but any suitable promoter can be used.

Preferably, recourse is made to a promoter regulation sequence which favours the overexpression of the coding sequence, such as, for example, that comprising at



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least one histone promoter such as described in European Application EP 0507698.

According to the invention, it is equally possible to use, in association with the promoter regulation sequence, other regulation sequences which are situated between the promoter and the coding sequence, such as "enhancer" transcription activators, such as, for example, tobacco etch virus (TEV) translation activator described in the Application WO87/07644, or of transit peptides, either single, or double, and in this case possibly separated by an intermediate sequence, that is to say comprising, in the transcription direction, a sequence coding for a transit peptide of a plant gene coding for a plastid localization enzyme, a part of the sequence of the N-terminal mature part of a plant gene coding for a plastid localization enzyme, then a sequence coding for a second transit peptide of a plant gene coding for a plastid localization enzyme, formed by a part of the sequence of the N-terminal mature part of a plant gene coding for a plastid localization enzyme, such as described in European Application No. 0 508 909. The invention provides a chimeric gene which comprises, between the promoter regulation sequence and the coding sequence, an optimized 25 transit peptide comprising, in the transcription sense, a sequence coding for a transit peptide of a plant gene coding for an enzyme with plastid localization, a part of the sequence of the N-terminal mature part of a plant gene coding for an enzyme



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with plastid localization, then a sequence coding for a second transit peptide of a plant gene coding for an enzyme with plastid localization.

It is possible to use as terminator or

polyadenylation regulation sequence any corresponding
sequence of bacterial origin, such as, for example, the
nos terminator of Agrobacterium tumefaciens, or even of
plant origin, such as, for example, a histone
terminator such as described in European Application



- 6 -

EP No. 0 633 317.

An object of the present invention is also plant cells, of monocotyledonous or dicotyledonous plants, especially of crops, transformed according to 5 one of the processes described above and comprising in their genome an efficacious quantity of a gene expressing hydroxyphenylpyruvate dioxygenase (HPPD). The invention also provides a plant regenerated from transformed plant cells. It has been observed that plants 10 transformed in this way have a significant tolerance to certain novel herbicides such as the isoxazoles described especially in French Patent Applications 9506800 and 95 13570 and especially of 4-[4-CF3-2-(methylsulphonyl)benzoyl]-5cyclopropylisoxazole, and especially isoxaflutole, a 15 selective maize herbicide, the diketonitriles such as those described in European Applications 0 496 630, 0 496 631, in particular 2-cyano-3-cyclopropyl-1-(2-SO₂CH₃-4-CF₃-phenyl)propane-1,3-dione and 2-cyano-3cyclopropyl-1-(2-SO₂CH₃-4-2,3-Cl₂-phenyl)propane-1,3-20 dione, the triketones described in European Applications 0 625 505 and 0 625 508, in particular sulcotrione.

method of weeding plants, especially crops, with the

25 aid of a herbicide of this type, characterized in that
this herbicide is applied to plants transformed
according to the invention, both pre-sowing, preemergence and post-emergence of the crop.

An object of the invention is also the use of

the HPPD gene as a marker gene in the course of the "transformation-regeneration" cycle of a plant species and selection on the above herbicide.

The different aspects of the invention will be better understood with the aid of the experimental examples below.

Example 1: Isolation of the HPPD gene of P. fluorescens A32

Starting from the amino-acid sequence of HPPD of Pseudomonas sp. P.J. 874 (published by Rüetschi U. et al., 1992, Eur. J. Biochem. 205: 459-466), the sequence of different oligonucleotides is deduced in order to amplify by PCR a part of the coding sequence of HPPD of P. fluorescens A 32 (isolated by McKellar,

- 15 R.C. 1982, J. Appl. Bacteriol., 53: 305-316). An amplification fragment of the gene of this HPPD has been used to screen a partial genomic bank of P. fluorescens A32 and thus to isolate the gene coding for this enzyme.
- 20 A) Preparation of genomic DNA of
 P. fluorescens A32

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The bacteria was cultivated in 40 ml of M63 minimum medium $(KH_2PO_4\ 13.6\ g/l,\ (NH_4)_2SO_4\ 2\ g/l,\ MgSO_4$ 0.2 g/l, FeSO₄ 0.005 g/l, pH 7 plus L-tyrosine 10 mM as the sole carbon source) at 28°C for 48 hours.

After washing, the cells are taken up in 1 ml of lysis buffer (100 mM tris HCl, pH 8.3, 1.4 M NaCl and 10 mM EDTA) and incubated for 10 minutes at 65°C.

After a phenol/chloroform treatment (24:1) and a chloroform treatment, the nucleic acids are precipitated by addition of one volume of isopropanol, then taken up in 300 μ l of sterile water and treated with final 10 μ g/ml RNAse. The DNA is treated afresh with phenol/chloroform, chloroform and reprecipitated by addition of 1/10 of the volume of 3 M sodium acetate, pH 5 and 2 volumes of ethanol. The DNA is then taken up in sterile water and determined.

B) Choice of the oligonucleotides and syntheses

Starting from the amino-acid sequence of HPPD of Pseudomonas sp. P.J. 874, five oligonucleotides are chosen, two directed in the terminal NH₂ direction of the protein towards the COOH terminal of the protein and three directed in the opposite direction (see Fig. 1). The choice was dictated by the two following rules:

- a stable 3' end of the
- oligonucleotide, that is to say at least two bases without ambiguity.
 - the smallest degeneracy possible.

The oligonucleotides chosen have the following sequences:

P1: 5'TA(C/T)GA(G/A)AA(C/T)CCIATGGG3'

P2: 5'GA(G/A)ACIGGICCIATGGA3'

P3: 5'AA(C/T)TGCATIA(G/A)(G/A)AA(C/T)TC(C/T)TC3'

P4: 5'AAIGCIAC(G/A)TG(C/T)TG(T/G/A)ATICC3'



P5: 5'GC(C/T)TT(A/G)AA(A/G)TTICC(C/T)TCIC3'

They were synthesized on a Cyclone plus DNA synthesizer of the make MILLIPORE.

With these five oligonucleotides, the

amplification fragments which must be obtained
theoretically by PCR starting from the sequence SEQ ID

No. 1 have the following sizes:

with the primers P1 and P3 ---->
approximately 690 bp

with the primers P1 and P4 ---->
approximately 720 bp

with the primers P1 and P5 ---->
approximately 1000 bp

with the primers P2 and P3 ---->

15 approximately 390 bp

with the primers P2 and P4 ---->
approximately 420 bp

with the primers P2 and P5 -----> approximately 700 bp

C) Amplification of a coding part of HPPD of P. fluorescens A32.

The amplifications were carried out on a PERKIN ELMER 9600 PCR apparatus and with PERKIN ELMER Taq polymerase with its buffer under standard conditions, that is to say for 50 μ l of reaction mixture there are dNTP at 200 μ M, primers at 20 μ M, 2.5 units of Taq polymerase and 2.5 μ g of DNA of

P. fluorescens A32.

The amplification programme used is 5 min at 95°C and then 35 <45 sec 95°C, 45 sec 49°C, 1 min 72°C> cycles followed by 5 min at 72°C.

Under these conditions, all the amplification fragments obtained have a size compatible with the theoretical sizes given above, which is a good indication of the specificity of the amplifications.

The amplification fragments obtained with the sets of primers Pl/P4, Pl/P5 and P2/P4 are ligated into pBSII SK(-) after digestion of this plasmid by Eco RV and treatment with the terminal transferase in the presence of ddTTP as described in HOLTON T.A. and GRAHAM M.W. 1991, N.A.R., Vol. 19, No. 5, p. 1156.

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A clone of each of the three types is

partially sequenced; this allows it to be confirmed
that a part of the coding region of the HPPD of

P. fluorescens A32 has indeed been amplified in the
three cases. The P1/P4 fragment is retained as probe in
order to screen a partial genomic bank of

20 P. fluorescens A32 and to isolate the complete gene of the HPPD.

D) Isolation of the gene

By Southern it is shown that a 7 Kbp fragment hybridizes, after digestion of the DNA of P.

25 fluorescens A32 by the restriction enzyme BamHI, with the probe HPPD P1/P4. 400 μg of DNA of P. fluorescens A32 are thus digested with the restriction enzyme BamHI and the DNA fragments making up approximately 7 Kbp are

purified on agarose gel.

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These fragments are ligated into pBSII SK(-), the latter is digested with BamHI and dephosphorylated by treatment with alkaline phosphatase. After transformation in E. coli DH10b, the partial genomic bank is screened with the probe HPPD P1/P4.

A positive clone was isolated and called pRP A. Its simplified map is given in Figure 2. On this map is indicated the position of the coding part of the HPPD gene. It is composed of 1077 nucleotides which code for 358 amino acids (see SEQ ID No. 1). The HPPD of P. fluorescens A32 has a good amino-acid homology with that of Pseudomonas sp. strain P.J. 874, in fact there is 92 % agreement between these two proteins (see Fig. 3).

Example 2: Construction of two chimeric genes

To confer plant tolerance to herbicides

inhibiting HPPD, two chimeric genes are constructed:

The first consists in putting the coding part
of the HPPD gene of P. fluorescens A32 under the
control of the double histone promoter (European Patent
No. 0 507 698) followed by tobacco etch virus
translational enhancer (TEV) (pRTL-GUS (Carrington and
Freed, 1990; J. Virol. 64: 1590-1597)) with the

terminator of the nopaline synthase gene. The HPPD will then be localized in the cytoplasm.

The second will be identical to the first, except that the optimized transit peptide (OTP) is



intercalated between the TEV transcription activator and the coding part of the HPPD (European Application EP No. 0 508 909). The HPPD will then be localized in the chloroplast.

- A) Construction of the vector pRPA-RD-153:
- pRDA-RD-11 A derivative of pBS-II SK(-)

 (Stratagene catalog #212206) containing the

 polyadenylation site of nopaline synthase (NOS polyA)

 (European Application EP No. 0 652 286) is cloned

 between the KpnI and SalI sites. The KpmI site is

 transformed into a NotI site by treatment with T4 DNA

 polymerase I in the presence of 150 μM of

 deoxynucleotide triphosphates and then ligation with an

 NotI linker (Stratagene catalog #1029). An NOS polyA

 cloning cassette is thus obtained.
 - pRPA-RD-127: A derivative of pRPA-BL-466
 (European Application EP No. 0 337 899) cloned in
 pRPA-RD-11 creating an expression cassette of the oxy
 gene and containing the promoter of the small subunit
 of ribulose biscarboxylase:

"promoter (SSU) - oxy gene - NOS polyA"

To create this plasmid, pRPA-BL-488 was
digested with XbaI and HindIII to isolate a fragment of
1.9 kbp comprising the SSU promoter and the oxy gene
which was ligated into the plasmid pRPA-RD-11, digested
with compatible enzymes.

- pRPA-RD-132: This is a derivative of pRPA-BL-488 (European Application EP No. 0 507 698) cloned



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into pRPA-RD-127 with creation of an expression cassette of the *oxy* gene with the double histone promoter:

"double histone promoter - oxy gene - NOS polyA"

To produce this plasmid, pRPA-BL-466 is
digested with HindIII, treated with Klenow and then
redigested with NcoI. The purified fragment of 1.35 kbp
containing the histone double promoter H3A748 is
ligated with the plasmid pRPA-RD-127 which had been
digested with XbaI, treated with Klenow and redigested
with NcoI.

- pRPA-RD-153: This is a derivative of
pRPA-RD-132 containing the translation activator of the
tobacco etch virus (TEV). pRTL-GUS (Carrington and
15 Freed, 1990; J. Virol. 64: 1590-1597) is digested with
NcoI and EcoRI and the 150 bp fragment is ligated into
pRPA-RD-132 digested with the same enzymes. An
expression cassette containing the promoter:
"double histone promoter - TEV - oxy gene - NOS polyA"
is thus created.

B) Construction of the vector pRPA-RD-185:

pUC19/GECA: A derivative of pUC-19 (Gibco
catalog #15364-011) containing numerous cloning sites.

pUC-19 is digested with EcoRI and ligated with the
oligonucleotide linker 1:

Linker 1: AATTGGGCCA GTCAGGCCGT TTAAACCCTA GGGGGCCCG

CCCGGT CAGTCCGGCA AATTTGGGAT CCCCCGGGC TTAA

The selected clone contains an EcoRI site



followed by the polylinker which contains the following sites: EcoRI, ApaI, AvrII, PmeI, SfiI, SacI, KpnI, SmaI, BamHI, XbaI, SalI, PstI, SphI and HindIII.

pRPA-RD-185: this is a derivative of

pUC19/GECA containing a modified polylinker. pUC19/GECA

is digested with HindIII and ligated with the

oligonucleotide linker 2:

Linker 2: AGCTTTAAT TAAGGCGCGC CCTCGAGCCT GGTTCAGGG

AAATTA ATTCCGCGCG GGAGCTCGGA CCAAGTCCC TCGA

The selected clone contains a HindIII site in the centre of the polylinker which now contains the following sites: EcoRI, ApaI, AvrII, PmeI, SfiI, SacI, KpnI, SmaI, BamHI, XbaI, SalI, PstI, SphI, HindIII, PacI, AscI, XhoI and EcoNI.

C) Construction of the vector pRP T:

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- pRP O: a derivative of pRPA-RD-153

 containing an expression cassette of HPPD, double

 histone promoter TEV HPPD gene terminator Nos. To

 produce pRP O, pRPA-RD153 is digested with HindIII,

 treated with Klenow and then redigested with NcoI to

 remove the oxy gene and replace it by the HPPD gene

 coming from the pRP A plasmid by BstEII digestion,

 Klenow treatment and redigestion with NcoI.
- pRP R: to obtain the plasmid, pRP O was
 digested with PvuII and SacI, the chimeric gene was
 purified and then ligated into pRPA-RD-185 and the
 latter was digested with PvuII and SacI.
 - pRP T: was obtained by ligation of the

chimeric gene coming from pRP R after digestion with SacI and HindIII in the plasmid pRPA-BL 150 alpha2 digested with the same enzymes (European Application EP No. 0 508 909).

The chimeric gene of the pRP T vector thus has the following structure:

| Double histone | TEV | Coding region | nos |
|----------------|-----|---------------|------------|
| promoter | | of HPPD | terminator |

D) Construction of the pRP V vector

- pRP P: this is a derivative of pRPA-RD-7

(European Application EP No. 0 652 286) containing the optimized transit peptide followed by the HPPD gene. It was obtained by ligation of the coding part of HPPD coming from pRP A by BstEII and NcoI digestion, Klenow treatment and from the plasmid pRPA-RD-7, the latter digested with SphI and AccI and treated with DNAse polymerase T4.

- pRP Q: a derivative of pRPA-RD-153

containing an expression cassette of HPPD, double

20 histone promoter - TEV - OTP - HPPD gene - Nos

terminator. To construct it, the plasmid pRPA-RD-153 is

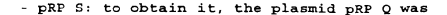
digested with SalI, treated with Klenow and then

redigested with NcoI to remove the oxy gene and replace

it by the HPPD gene released from the pRP P plasmid by

BstEII digestion, Klenow treatment and redigestion with

NcoI.





digested with PvuII and SacI to release the chimeric gene, which was ligated into pRPA-RD-185, the latter digested with PvuII and SacI.

- pRP V: it was obtained by ligation of the chimeric gene released from pRP S, after digestion with SacI and HindIII, into the plasmid pRPA-BL 150 alpha2 (European Application EP No. 0 508 909).

The chimeric gene of the pRP Q vector thus has the following structure:

| 10 | Double histone | TEV | OTP | Coding region | nos |
|----|----------------|-----|-----|---------------|------------|
| | promoter | | | of HPPD | terminator |

Example 3: Transformation of the industrial tobacco PBD6

In order to determine the efficacy of these
two chimeric genes, these were transferred to
industrial tobacco PBD6 according to the transformation
and regeneration procedures already described in
European Application EP No. 0 508 909.

1) Transformation:

The vector is introduced into the nononcogenic strain of Agrobacterium EHA 101 (Hood et al.,
1987) which carries the cosmid pTVK 291 (Komari et al.,
1986). The transformation technique is based on the
procedure of Horsh R. et al. (1985), Science, 227,

2) Regeneration:

1229-1231.

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The regeneration of the tobacco PBD6 (origin

SEITA France) from foliar explants is carried out on a Murashige and Skoog (MS) base medium comprising 30 q/l of sucrose as well as 200 ug/ml of kanamycin. The foliar explants are selected on plants in the greenhouse or in vitro and transformed according to the foliar discs technique (Science 1985, Vol. 227, p. 1229-1231) in three successive steps: the first comprises the induction of shoots on an MS medium to which is added 30 g/l of sucrose containing 0.05 mg/l 10 of naphthylacetic acid (NAA) and 2 mg/l of benzylaminopurine (BAP) for 15 days. The shoots formed during this step are then developed by culture on an MS medium to which is added 30 g/l of sucrose, but not containing any hormone, for 10 days. Developed shoots 15 are then selected and cultured on an MS rooting medium of half salts, vitamins and sugars content and not containing any hormone. At the end of approximately 15 days, the rooted shoots are placed in earth. Example 4: Measurement of the tolerance of the tobacco to 4-[4-CF₃-2-(methylsulphonyl)benzoyl]-5-20 cyclopropylisoxazole: post-emergence treatment

On leaving in-vitro culture, the transformed tobacco plantlets were acclimatized in a greenhouse (60 % relative humidity; temperature: 20°C during the night and 23°C during the day) for five weeks and then treated with 4-[4-CF₃-2-(methylsulphonyl)benzoyl]-5-cyclopropylisoxazole.

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The control tobacco, non-transformed and

treated with 4-[4-CF₃-2-(methylsulphonyl)benzoyl]-5-cyclopropylisoxazole in doses ranging from 50 to 400 g/ha, develops chloroses in approximately 72 hours, which intensify to develop into very pronounced necroses in a week (covering approximately 80 % of the terminal leaves).

After transformation, this same tobacco, which overexpresses the HPPD of P. fluorescens, is very well protected against treatment with 4-[4-CF₃-2-(methylsulphonyl)benzoyl]-5-cyclopropylisoxazole at a dose of 400 g/ha.

If the overexpressed enzyme is in the cytoplasm, that is to say if the transformation was carried out with the gene carried by the vector pRP T, then the plant shows very slight chloroses which are all localized on the intermediate leaves.

If the overexpressed enzyme is in the chloroplast, that is to say if the transformation was carried out with the gene carried by the vector pRP V, then the plant is perfectly protected and does not show any symptoms.

Example 5: Measurement of the tolerance of the tobacco
to 4-[4-CF3-2-(methylsulphoyl)benzoyl]-5cyclopropylisoxazole: pre-emergence treatment

a) in vitro test:

Tobacco seeds harvested from plants from the "transformation-regeneration" cycle and resistant to isoxaflutole foliar treatment are used at a dose of



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400 g/h described in Examples 1 to 3.

These seeds were sown in boxes containing plant agar at 10 g/l and isoxaflutole at different concentrations ranging from 0 to 1 mg/l. Germination was then carried out at 25°C with a photoperiod of 12 hours of light/12 hours of darkness.

According to this protocol, wild tobacco seeds were germinated as well as seeds of the two types of transgenic tobacco, that is to say CY tobaccos, with localization of the HPPD in the cytoplasm, and the CO tobaccos with localization of the HPPD in chloroplast.

Resistance measurements are carried out visually between 2 and 3 weeks after sowing.

The results are recorded in the table below.

| isoxaflutole | Wild tobacco | CY tobacco | CO tobacco |
|--------------|------------------------|--|---|
| concentra- | | | |
| tion | | | |
| 0 mg/l | 100 % of the | 100 % of the | 100 % of the |
| | seeds ger- | seeds | seeds |
| | minate | germinate | germinate |
| | without | without | without |
| | symptoms° | symptoms° | symptoms |
| 0.05 mg/l | 20 % of the | 75 % of the | 75 % of the |
| | seeds | seeds | seeds |
| | germinate | germinate* | germinate* |
| | and show | without | without |
| | symptoms° | symptoms° | symptoms° |
| | concentra- tion 0 mg/l | concentra- tion 0 mg/l 100 % of the seeds ger- minate without symptoms° 0.05 mg/l 20 % of the seeds germinate and show | concentration 0 mg/l 100 % of the 100 % of the seeds ger-seeds minate germinate without symptoms° symptoms° 0.05 mg/l 20 % of the 75 % of the seeds germinate germinate* and show without |



| 0.1 mg/l | no | 75 % of the | 75 % of the |
|----------|-------------|-------------|-------------|
| | germination | seeds | seeds |
| | | germinate* | germinate* |
| | | without | without |
| | | symptoms° | symptoms° |
| 0.5 mg/l | no | 75 % of the | 75 % of the |
| | germination | seeds | seeds |
| | | germinate* | germinate* |
| | | without | without |
| | | symptoms° | symptoms° |
| 1 mg/1 | no | 75 % of the | 75 % of the |
| | germination | seeds | seeds |
| | | germinate* | germinate* |
| | | with slight | without |
| | | symptoms° | symptoms° |

- of germination are more or less significant deformations of the cotyledons and above all a bleaching of the tissues which are normally photosynthetic and thus green.
- * 75 % of the seeds germinate because seeds from the

 self-fertilization of single-locus plants coming from
 the "transformation-regeneration" cycle and thus only
 carrying the tolerance gene on one chromosome were
 sown.

Working in the same way with the following products,



Product No. 51 of American Patent 4 780 127, the same results are obtained at a concentration of 0 mg/l and 0.1 mg/l on wild tobacco and CO tobacco.

b) greenhouse test:

Measurement is carried out as in Example 4, apart from the treatment being carried out preemergence, 24 hours before sowing. Wild sowing is carried out normally. Under these conditions, it is observed that, for the non-treated control sowings, there is no germination for any dose of herbicide at least equal to 10 g/ha. On the contrary, the CY tobaccos do not show any symptoms, such as defined in paragraph a), up to and including 100 g/ha. Similarly, the CO tobaccos do not show any symptoms, such as defined in paragraph a), up to and including 200 g/ha.

These results show clearly that the HPPD gene of P. fluorescens confers a tolerance to the tobacco against pre-emergence treatments with isoxaflutole. This tolerance is better if the protein is localized in the chloroplast in place of the cytoplasm.

Example 6:

With the aim of studying whether the HPPD gene of Pseudomonas fluorescens can be used as a marker gene in the course of the "transformation-regeneration" cycle of a plant species, tobacco was transformed with the HPPD gene and transformed plants were obtained after selection on isoxaflutole.

Material and methods and results



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The chimeric gene pRP V described below is transferred into industrial tobacco PBD6 according to the transformation and regeneration procedures already described in European Application EP No. 0 508 909.

The chimeric gene of the vector pRP V has the following structure:

| Double histone | TEV | OTP | Coding region | nos |
|----------------|-----|-----|---------------|------------|
| promoter | | | of HPPD | terminator |

1) Transformation:

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The vector is introduced into the

Agrobacterium EHA 101 non-oncogenic strain (Hood et
al., 1987) which carries the cosmid pTVK 291 (Komari et
al., 1986). The transformation technique is based on
the procedure of Horsh et al. (1985).

15 2) Regeneration:

The regeneration of the tobacco PBD6 (origin SEITA France) from foliar explants is carried out on a Murashige and Skoog (MS) base medium comprising 30 g/l of sucrose as well as 350 mg/l of cefotaxime and 1 mg/l of isoxaflutole. The foliar explants are selected on plants in a greenhouse or in vitro and transformed according to the foliar discs technique (Science 1985, Vol. 227, p. 1229-1231) in three successive steps: the first comprises the induction of shoots on an MS medium to which is added 30 g/l of sucrose containing 0.05 mg/l of naphthylacetic acid (NAA) and 2 mg/l of benzylaminopurine (BAP) for 15 days and 1 mg/l of



isoxaflutole. The green shoots formed in the course of this step are then developed by culture on an MS medium to which are added 30 g/l of sucrose and 1 mg/l of isoxaflutole, but not containing hormone, for 10 days.

- Developed shoots are then selected and are cultured on an MS rooting medium of half salts, vitamins and sugars content and 1 mg/l of isoxaflutole and not containing any hormone. At the end of approximately 15 days, the rooted shoots are placed in earth.
- All the plantlets obtained according to this protocol are analysed by PCR with specific primers of the HPPD of P. fluorescens. This PCR analysis has enabled it to be confirmed that all the plantlets thus obtained have indeed integrated the HPPD gene.
- In conclusion, this assay confirms that the HPPD gene can be used as marker gene and that, associated with this gene, isoxaflutole can be a good selection agent.

Examples 7 and 8: Isolation of the HPPD gene of

Arabidopsis thaliana and of the HPPD gene of carrot

(Daucus carotta)

a) Construction of cDNA banks

mRNAs extracted from young plantlets of Arabidopsis

thaliana and mRNAs extracted from carrot cells in

culture served to construct two cDNA banks in the

vector Uni ZapTM XR marketed by the company Stratagen,

following the protocol recommended by this company.

b) Screening of the cDNA banks

These two banks were screened with the aid of a probe corresponding to a cDNA of Arabidopsis thaliana of partial length, obtained via the Arabidopsis Biological Resource Center (Ohio, USA) and indexed: EST clone No. 91B13T7. This clone is formed of approximately 500 base pairs of which only 228 had been sequenced by the MSU-DOE Plant Research Laboratory in the context of random sequencing of cDNA of Arabidopsis thaliana. We completely sequenced the 500 base pairs before using this clone to screen our cDNA banks of Arabidopsis thaliana and of carrot with the aid of the classical technique of hybridization of lysis regions (reference?).

- c) A cDNA of Arabidopsis thaliana (SEQ ID No. 2)

 15 corresponding to 1338 base pairs was obtained. This cDNA has a translation initiation start codon in position 25 and a translation end codon in position 1336. This cDNA is thus complete and codes for a protein of 445 amino acids.
- d) A cDNA of carrot (Daucus carotta) (SEQ ID No. 3) corresponding to 1329 base pairs was obtained. This cDNA has a translation initiation start codon in position 1 and a translation finish codon in position 1329. This cDNA is thus complete and codes for a protein of 442 amino acids.

The sequences illustrated are the following: SEQ ID No. 1 Sequence of the HPPD gene of Pseudomonas fluorescens A32



SEQ ID No. 2

cDNA sequence of HPPD of Arabidopsis thaliana SEO ID No. 3

cDNA sequence of HPPD of Daucus carotta

5 The figures below are given by way of indication to illustrate the invention.

Figure 1 represents the protein sequence of the HPPD of Pseudomonas sp. strain P.J 874 and the theoretical nucleotide sequence of the corresponding coding part;

the five oligonucleotides chosen to carry out the amplification of a part of this coding region are symbolized by the five arrows.

Figure 2 represents the mapping of the plasmid with the genomic DNA fragment of 7 kb comprising the gene of the

15 HPPD of P. fluorescens A32.

Figure 3 gives the comparison of the amino-acid sequences of the HPPD of P fluorescens A32 and of the HPPD of Pseudomonas sp. strain P.J.874 (only the divergent amino acids between the two sequences are indicated) as well as the concensus sequence.

Throughout this specification, unless the context requires otherwise, the word "comprise", or variations such as "comprises" or "comprising", will be understood to imply the inclusion of a stated element or integer or group of elements or integers but not the exclusion of any other element or integer or group of elements or integer.



Sequence listing

(1) GENERAL INFORMATION

- (i) APPL!CANT: Sailland, Alain Rolland, Anne Matringe, Michel Pallett Kenneth E
- (i) TITLE OF INVENTION: DNA SEQUENCE OF A HYDROXYPHENYLPYRUVATE DIOXYGENASE GENE AND OBTAINMENT OF PLANTS COMPRISING THIS HYDROXYPHENYLPYRUVATE DIOXYGENASE GENE, WHICH ARE RESISTANT TO CERTAIN HERBICIDES
- (iii) NUMBER OF SEQUENCES: 3
- (IV) CORRESPONDENCE ADDRESS:
 - (A) ADDRESSEE: Francois Chretien
 - (B) STREET: 14-20 rue Pierre BAIZET
 - (C) CITY: Lyon Cedex 09
 - (E) COUNTRY: France
 - (F) ZIP: 69263
- (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOSMS-DOS
 - (D) SOFTWARE: Patentin Release #1.0, Version #1.25
- (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER: FR PH95033
 - (B) FILING DATE: 02-JUN-1995
 - (C) CLASSIFICATION:
- (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: Chretien, Francois
- (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: 72-29-26-42
 - (B) TELEFAX: 72-29-28-43
- (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1077 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (genomic)
- (iii) HYPOTHETICAL: NO
- (N) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Pseudomonas fluorescens



(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1.

| 60 | CONNTINGCO | TIGATICAT | CTGATGGGCT | CCCANTESEC | TATACGAAAA | ATGGERGATE |
|------|-------------|------------|-------------|--------------|------------|--------------------------|
| :20 | EXAMPTEGES | TARRETTERS | TTOGAGATEA | GGAGGGGATT | 1530thccst | ********* |
| 130 | CATCOTCAAC | AGAICAACCT | CCCAGGGCG | GCACCTGTAC | ********** | CONTRACTOR |
| 240 | SCTCTSCGGC | ACCCCCCCTC | GCGGCCGAAC | CICCIACITI | ACAGCATOGO | ಸ್ವಾ ಚಿ |
| 300 | ACTCGGCGCC | CCCCCTCCA | GCCTACAACC | CTCGCANAG | SCSTGAAGGA | ATGGCSTTCC |
| 360 | CAAGGGCATC | TGCCGGCGAI | GAATTGAACC | COGGCCGATG | ATATTGACAC | عموده |
| 420 | CTACGACATC | GCAGCTCGAT | TTCGGCGAAG | CATCGACCGT | CGTTGTACCT | ಯಲಾಯಾಯ |
| 480 | CANASTONIC | GTGCAGGTCT | MICCEGTCG | TOTGGAGCGC | ACCTCGAAGG | ೧೩೦೦೦ |
| 540 | CTTCTACGAG | ACTGGGCCAA | CCCATCCTCT | CTATCGCGGC | CCCACAACGT | CACCACCTCA |
| 600 | CACCGGCCTG | AGGGCGAGTA | TTCGATATCA | AGCGCGTTAC | ACTTCCCTCA | AMITOTICA |
| 660 | CONGACTCG | TCCCGCTGAA | ATGATCCGCA | CCCGGACGGC | CCATGAGTGC | ACTTCCAAGG |
| 720 | AGGCATCCAG | TCAACGGCGA | CTCATGCAGT | CCANGAGTTC | CGGGGCACAT | TOCONSCIOCO |
| 760 | GANANTCGGC | ACCCUTTONA | AAGACCTGGG | CGASCTGGTC | TCCTCACCGA | CACGTGGCGT |
| 840 | CONCETTOCCT | TOCTCGAAGG | TATTACGANA | GCCAGAGACACT | TGACCGCGCC | A:GCGCTTCA |
| 900 | COGATOTTCC | TCCTGCTGGA | GENEGGGEN | TCAACTECAG | AGCCGGTGGA | فمحجمحهج |
| 960 | GGGCCCGGTG | AMECUTEAT | ATCTTCTCGG | GCTGCTGCAG | ACANACGCCT | g . ccauseccs |
| 1020 | CTTCAMEGCG | CCENCCCCIA | GAIGGGITTG | CANGGGCGAC | TCATCCAGCG | TETTCENI |
| 1077 | CGATTAA | TATTGACCGC | carcaracara | TGACCAGGTG | CCATCGAACG | CTCTTCCACT |
| | | | | | | |

(2) INFORMATION FOR SEQ ID NO.2:

- (1) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1338 base pairs

 - (B) TYPE: nucleic scid (C) STRANDEDNESS: double (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE:
 - (A) ORGANISM: Arabidopsis thaliana (B) STRAIN: Columbia

 - (D) DEVELOPMENTAL STAGE: Young green plant
- (vii) IMMEDIATE SOURCE:
 (A) LIBRARY: Uni zap XR STRATAGENE
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:
- ATGGGGGGACC AAAAGGGGGG CGTTTCAGAG AATGAAAACC ATGATGACGG CGCTGCGTCG 60 TOGGCOGGAI TOMAGTOGT COGATITICS MAGTTOGTAA GUAGAATCC MAGTCTGAT 120



| WITCHES | TTAAGCGCTT | CCATCACATO | CACTICICS | COCCOCCACCC | MICANCUTC | 18 |
|---|---------------|--------------|------------|-------------|------------|------|
| STT:ST:SC: | TOTOCTOCCO | TETEGGGGATG | AGATTCTCCC | CCAAATCCCA | TCTTTCCACC | 24 |
| ;;;kaakaatsa | TEACGESTS | בדבאנדבבאנדב | ACCTCCGGTG | ACCTCEGATT | SUTTEMET | 30 |
| ; ;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;;; | वाध्यक्षात्रस | enscences ex | CAGATTANAC | CONCURCENC | AUCTICIAIC | 36 |
| COTTTOAKTE | ATTACOGCTC | Herente | 11C11C1C11 | CACAISGICT | COSTOTTAGA | 42 |
| sco ort cca | TTGAAGTAGA | AGACGCAGAG | TCAGCTTTCT | CCATCAGTGT | AGCTAATGGC | 48 |
| SCIATICUTE | CCICCCCICC | TATCGTCCTC | MICHAGOAG | TTACCATCCC | TOAGGTTANA | 54 |
| CTATACGGCG | AIGTIGITOT | CCGATATGTT | AGTTACANAG | CACHIGATAC | COMMITTEE | 60 |
| صيتصنو | CAGGGTTCGA | CCCTCTACAG | CATCCCTCCT | COTTCCCATT | CONTINICCT | 66 |
| YECCESCEEC | TTGACCACGC | COTGGGGGGG | CTTCCTCASC | TTCCTCCCCC | TTIANCTTAT | 720 |
| CTAGCGGGGT | TCACTGGTTT | TCACCUATTC | GCAGAGTTCA | CAGCAGACGA | CGTTGGNACC | 780 |
| eccenenece | GTTTANATTC | AGCGGTCCTG | GCTAGCAAIG | atguatggt | TCTTCTACCS | 840 |
| ATTAACGAGC | CACTGCACGG | YYCYYYCYCC | AAGAGTCAGA | TTOMOLOGIA | TTTCGAACAT | 900 |
| MCGMGGCG | CAGGGCTACA | ACAICTESCT | CTGATGAGTG | ANGACATATT | CAGGACCCTG | 960 |
| ACAGAGATGA | CONTINUENT | CAGTATTOGA | CONTICGACT | TCATCCCTTC | TCCTCCGCCT | 1020 |
| ACTIACTACE | ACANICICAN | CANACCECTE | GGCGACGTGC | TCAGCGATGA | TOWERTONS | 1080 |
| CACTGTCAGG | AASTAGGGAI | TOTTGTAGAC | AGAGATGAIC | AAGGGACGTT | SCITCALLIC | 1140 |
| TTCACAAAAC | CACTAGGTGA | cyceccance | ATATTTATAG | AGATAATCCA | CACACTACCA | 1200 |
| TGCATGATGA | AAGATGAGGA | AGGGAAGGCT | TACCAGAGTG | CACCATCTCC | TECTTTTESC | 1260 |
| ANAGGENATT | TCTCTGAGCT | CITCANGICC | ATTONOUT | remmere | TCTTGANGCC | 1320 |
| ALACAGTTAG | TOGGATCA | | | | | 1338 |

(2) INFORMATION FOR SEQ ID NO:3:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1329 base pairs

 - (B) TYPE: nucleic acid
 (C) STRANDEDNESS: double
 (D) TOPOLOGY: Enear
- (ii) MOLECULE TYPE: cONA
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO
- (vi) ORIGINAL SOURCE: (A) ORGANISM: Daucus carota
 - (D) DEVELOPMENTAL STAGE: Suspension cells
- (vii) IMMEDIATE SOURCE:
 - (A) LIBRARY: Uni zap XR STRATAGENE
- (x) SEQUENCE DESCRIPTION: SEQ ID NO:3:

ATGGGGAAAA AACAATGGGA AGCTGAAATT CTCTCAAGCA ATTCATCAAA CACCTCTCCT



| DIALIATTEA | AGETGGTSSG | TTTEAREAGE | TTEGTSEGES | SENYSSERY | GTOCGATCAC | 120 |
|----------------------|-------------------|------------|------------|------------|------------|------|
| tttscccetex | AGCGGTTCCA | CCACATTGAS | TTCTGGTGCG | GCGACGCCAC | CAACACGTCG | 130 |
| CSGCGGTTGT | CGTGGGGCCT | COCCATOCCT | TECTCCCCY | ANTIGGNICT | CTCTACTGGA | 240 |
| AACTCTGTTC | ACGCTTCTTA | TCTTGTTCGC | TOGGOGAATO | TCAGTTTCGT | CTTCACCGCT | 300 |
| CCTTACTCTC | CCTCCACCAC | CACTTCCTCT | GCTTCAGCTG | CCATCCCGTC | TTTTTCGGCA | 360 |
| TCGGGTTTTC | ACTOTTTTGC | GGCCANACAC | GCCCTTGCTG | TICGGGCTAI | TGCTCTTGAA | 420 |
| CTTCCTCACG | TGGCTGCTGC | GTTTGAGGCC | ACTGTTGCCC | GTGGGGCCAG | SCCGGCTTCG | 480 |
| GCTCCTGTTG | AATTGGACGA | CCAGGCGTGG | TTGGCTGAGG | TGGAGTTGTA | CGGAGATGTG | 540 |
| CICTICALCI | TIGTIAGTII | TGGGAGGGAG | CAGGGTTTGT | TITICCIES | ATTCGAGGCG | 600 |
| CTCCACCCCA | CGGCGTCGTT | TCCGGATTTG | GATTATGGAA | TTAGANGACT | TGATCATGCG | 560 |
| GTGGGGAATG | TTACCGAGTT | GGGGCCTGTG | GTGGAGTATA | TTANAGGGTT | TACGGGGTTT | 720 |
| CATCUATTTG | CCCACTITAC | AGCGGAGGAI | GTGGGGACTT | TGGAGAGTGG | CTTCAATTCG | 780 |
| ः ब्लाब्स द्र | CCAATAATCA | GGAGATGGTT | CTGTTGCCCT | TGANTENGCC | TGTGTATGGG | 840 |
| ACCANGAGGA | AGACTCAGAT | ACAGACTIAC | TTGGAGCACA | ATCHAGGGGC | TGGAGTGCAG | 900 |
| CATTTGGCTT | TAGTGAGTGA | GGATATTTT | AGGACTITAA | GCCACATCAG | GAAGAGGAGT | 960 |
| recerreere | GTTTTGAGTT | TATGCCTTCG | CCACCGCCTA | CCTATTACAA | CAATTTCAAG | 1020 |
| AATAGGGTCG | GGGATGTGTT | CACTCATGAX | CAGATCAAGG | AGTGTGAAGA | TTTGGGGATT | 1080 |
| TTGGTGGATA | GGGATGATCA | GGGTACATTG | CTTCAAATCT | TTACCAAGCC | TGTAGGTGAC | 1140 |
| ASSCCTACCT | TATTCATAGA | CATCATTCAG | AGGGTAGGGT | GCATGCTCAA | GGACGATGCA | 1200 |
| TOTACACOOS | ACCAGAAGGG | CGGGTGCGGA | CCATTTCCCA | AGGGGAACTT | CTCAGAGCTG | 1260 |
| TTCANGTCCA | TCGAAGAATA | TGAAAAAA | CTTGAAGCTA | AACAAATCAC | TGGATCTGCT | 1320 |
| CTSCATGA | | | | | | 1329 |



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THE CLAIMS DEFINING THE INVENTION ARE AS FOLLOWS:

- 1. Isolated sequence of a gene expressing a hydroxyphenylpyruvate dioxygenase (HPPD) for producing 5 transgenic plants that are tolerant to certain herbicides, characterised in that said sequence is obtainable from Pseudomonas fluorescens and comprises the sequence set forth in SEQ ID NO:1.
- 10 2. Isolated sequence of a gene expressing a hydroxyphenylpyruvate dioxygenase (HPPD) for producing transgenic plants that are tolerant to certain herbicides, characterised in that said sequence is of plant origin.
- 15 3. Sequence according to claim 2, characterized in that it is from Arabidopsis.
 - 4. Sequence according to claim 3, comprising the sequence set forth in SEQ ID NO:2.
 - 5. Sequence according to claim 2, characterized in that it is from an umbelliferous plant.
- 6. Sequence according to claim 5, wherein the 25 umbelliferous plant is a carrot.
 - 7. Sequence according to claim 6, comprising the sequence set forth in SEQ ID NO:3.
- 30 8. Chimeric gene for the genetic transformation of plants comprising, in the transcription sense:
 - at least one promoter regulation sequence from a gene expressed naturally in plants,
 - a heterologous coding sequence,
- 35 at least one polyadenylation sequence, characterized in that the coding sequence is a sequence of a gene which expresses

- 31 -

a hydroxyphenylpyruvate dioxygenase (HPPD).

- 9. Chimeric gene according to claim 8, characterized in that the coding sequence is of bacterial or of plant origin.
- 10. Chimeric gene according to claim 9, characterized in that the coding sequence is from Pseudomonas sp.
- 11. Chimeric gene according to claim 8, characterized in 10 that the coding sequence is a sequence such as defined in any one of claims 1 to 7.
- 12. Chimeric gene according to any one of claims 9 to 11, characterized in that the promoter regulation sequence favours 15 the overexpression of the coding sequence.
 - 13. Chimeric gene according to claim 12, characterized in that the promoter regulation sequence comprises at least one histone promoter.
 - 14. Chimeric gene according to any one of claims 9 to 13, characterized in that it comprises a transit peptide between the promoter regulation sequence and the coding sequence.
- 25 15. Chimeric gene according to any one of claims 9 to 14, characterized in that it comprises, between the promoter regulation sequence and the coding sequence, an optimized transit peptide comprising, in the transcription sense, a sequence coding for a transit peptide of a plant gene coding for an enzyme with plastid localization, a part of the sequence of the N-terminal mature part of a plant gene coding for an enzyme with plastid localization, then a sequence coding for a second transit peptide of a plant gene coding for an enzyme with plastid localization.



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- 16. Chimeric gene according to any one of claims 9 to 15, characterized in that it comprises a sequence of a transcription activator (enhancer) between the promoter regulation sequence and the coding sequence.
- 17. Vector utilizable for the genetic transformation of plants, characterized in that it comprises a chimeric gene according to any one of claims 9 to 16.
- 18. Plant cell, characterized in that it comprises a chimeric gene according to any one of claims 9 to 16.
 - 19. Plant, characterized in that it is regenerated from cells according to claim 18.
 - 20. Plant according to claim 19, characterized in that it belongs to the dicotyledonous plant family.
- 21. Plant according to claim 20, characterized in that it 20 is a tobacco plant.
 - 22. Plant according to claim 21, characterized in that it is tobacco PBD6.
- 23. Process of transformation of plants to make them tolerant to inhibitors of HPPD, characterized in that a gene expressing an exogenous HPPD is introduced into a vegetable cell.
- 30 24. Process according to claim 23, characterized in that the transfer is carried out with Agrobacterium tumefaciens or Agrobacterium rhizogenes.
- 25. Process according to claim 23, characterized in that 35 the transfer is carried out by delivery by bombardment with the aid of charged particles of the DNA.



- 26. Process of transformation of plants, characterized in that a gene expressing an exogenous HPPD is introduced into the vegetable cell as a selection marker.
- 27. Process according to any one of claims 23 to 26, characterized in that a chimeric gene according to any one of claims 8 to 16 is introduced.
- 28. Process of selective herbicide treatment of plants, 10 characterized in that an HPPD inhibitor is applied to a transformed plant comprising cells according to claim 18.
- 29. Process according to claim 28, characterized in that the plants are the plants according to any one of claims 19 to 15 22.
 - 30. Process according to any one of claims 28 and 29, characterized in that the plants are obtained by the process according to claims 23 to 27.
 - 31. Process according to any one of claims 28 to 30, characterized in that the HPPD inhibitor is an isoxazole, a diketonitrile, a triketone or sulcotrione.
- 25 32. Process according to claim 31, characterized in that the isoxazole is 4-[4-CF₃-2-(methylsulphonyl)benzoyl]-5-cyclopropylisoxazole.
- 33. Sequence according to any one of claims 1 to 7, 30 substantially as described herein with reference to the Figures and/or Examples.
- 34. Chimeric gene according to any one of claims 8 to 16, substantially as described herein with reference to the Figures 35 and/or Examples.



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- 34 -

- 35. Vector according to claim 17, substantially as described herein with reference to the Figures and/or Examples.
- 36. Plant cell according to claim 18, substantially as 5 described herein with reference to the Figures and/or Examples.
 - 37. Plant according to any one of claims 19 to 22, substantially as described herein with reference to the Figures and/or Examples.

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38. Process according to any one of claims 23 to 32, substantially as described herein with reference to the Figures and/or Examples.

DATED this FIRST day of MARCH 2000 Rhone-Poulenc Agro

By DAVIES COLLISON CAVE Patent Attorneys for the Applicants



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Fig 1

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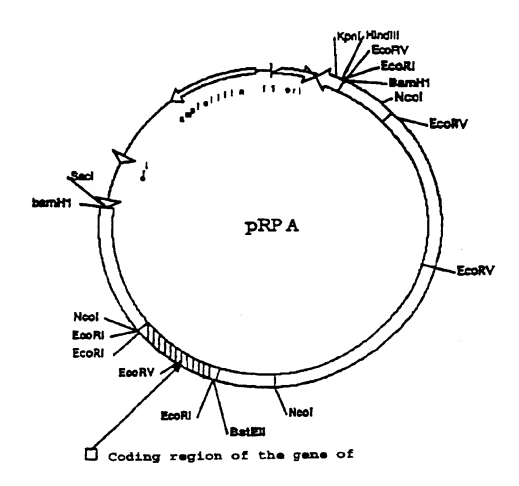


Fig 2.

2/3

| Consensus | | | SPTP.TLEPI | | | Se |
|-----------------------------------|------------|------------|------------|------------|-------------|------------------------|
| P. fluorescens Pseudomonas sp. | H | | G | ••••••• | N | 54 49 |
| Consensus | RQG.INLILH | NEP.S.ASYF | AAEHGPSVCG | MAFRYKDSQK | AY. RALELGA | 100 |
| P. fluorescens Pseudomonas sp. | E | H.V | •••••• | ******** | x | 1 00 99 |
| Consensus | QPIHI.TGPM | ELNLPAIKGI | GGAPLYLIDR | FGEGSSTYDI | DFV.LEGV.R | 150 |
| P. fluorescens Pseudomonas sp. | D | •••••• | •••••• | ********* | YE. | 150 149 |
| Consensus | .PVGAGLK.I | DHLTHNYYRG | RH.YWANFYE | KLFNFRE.RY | FDIXGEYTGL | 299 |
| P. fluorescens Pseudomonas sp. | NV. HI. | ••••• | v | A | ••••• | 200 199 |
| Consensus | TSKAH.APDG | MIRIPLNEES | SKGAGQIEEF | LNQFNGEGIQ | HVAFL.DDL. | 250 |
| P. fluorescens Pseudomonas sp. | S | | | | | 250 249 |
| Consensus | KTWD.LK.IG | NRFWTAPPOT | YYENLEGRLP | .HGEPYLQ | ARGILLDGSS | 300 |
| P. fluorescens Pseudomonas sp. | HS | | | | | 300 299 |
| Consensus | GDKRLLLQ | IFSETLNGPY | FFEFIQRKGD | DGFGEGNFKA | LFESIERDQV | 350 |
| P. fluorescens Pseudomonas sp. | VE ES | | | | | 35 0 349 |
| Consensus | RRGVLD | | | | | 358 |
| . fluorescens Seudomonas en. | TÀ. | | | | | 358 357 |

Fig. 3